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- GB 1571629

 - GB 1500613 GB 1500612 GB 1500611

 - GB 1520246
 - **GB 147946** GB 1478020
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- **Applicants**
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(54) Inhalation drugs

(57) A finely divided inhalation drug, e.g. sodium cromoglycate, comprises a therapeutically effective proportion of individual particles capable of penetrating deep into the lung, characterised in that a bulk of the particles which is both unagglomerated and unmixed with a coarse carrier, is sufficiently free flowing to be filled into capsules on an automatic filling machine and to empty from an opened capsule in an inhalation device. A proportion of the individual drug particles may have a spherical, collapsed spherical or ring dough-

Mr 6000

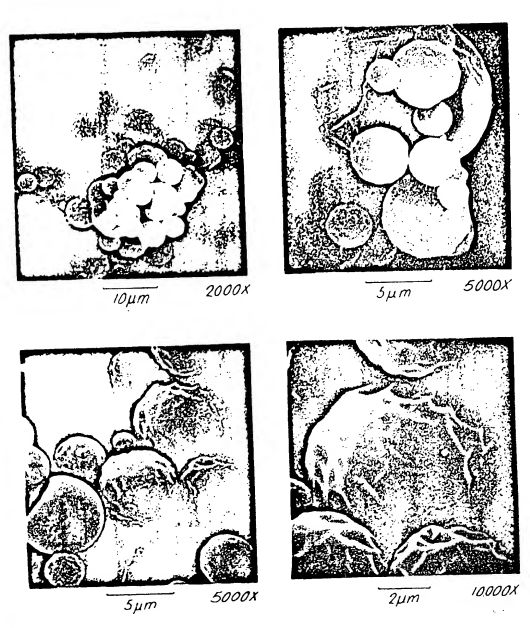


Fig.1.

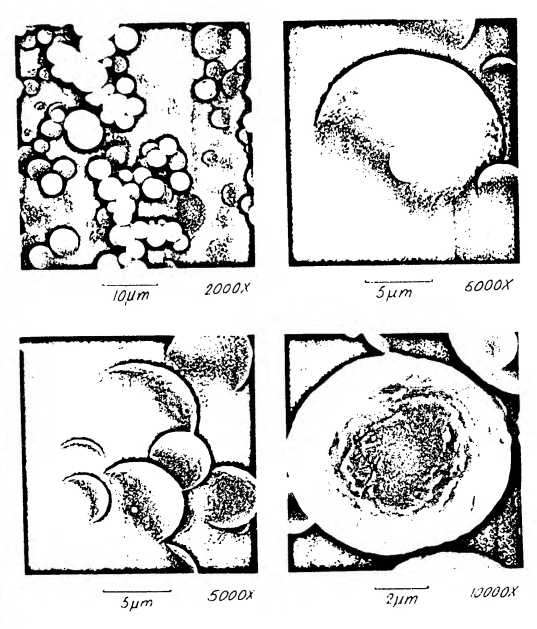


Fig.2.

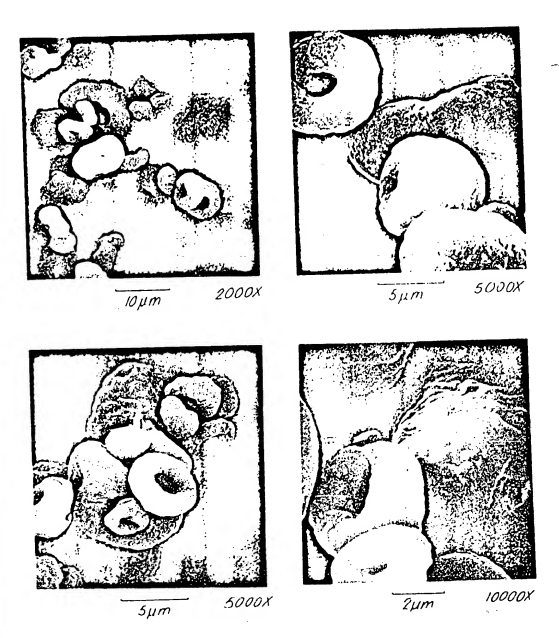


Fig. 3.

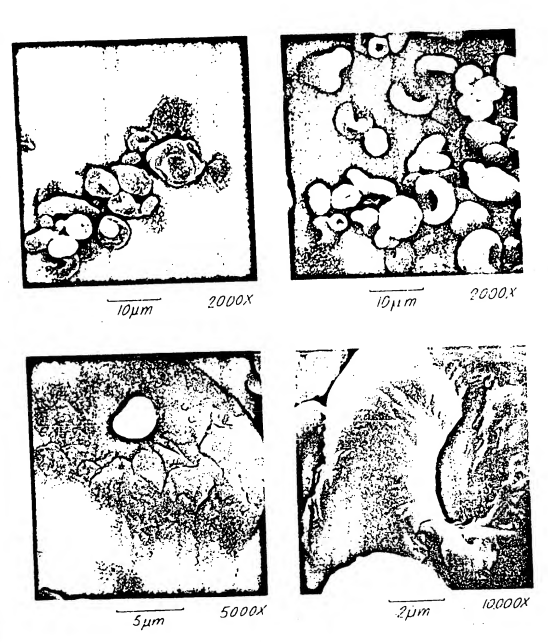


Fig.4.

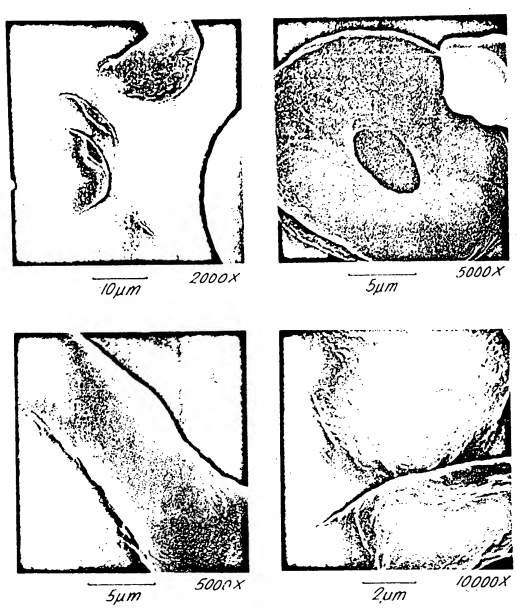


Fig. 5.

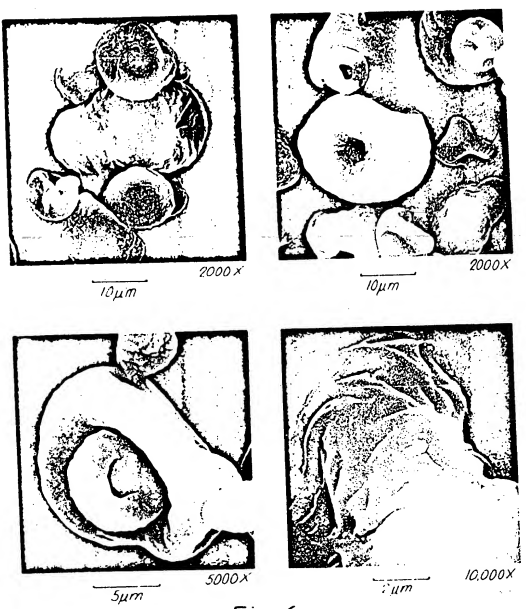
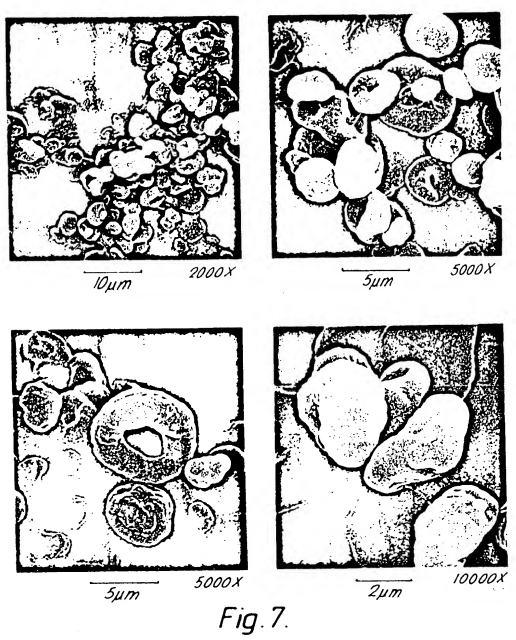


Fig.6.



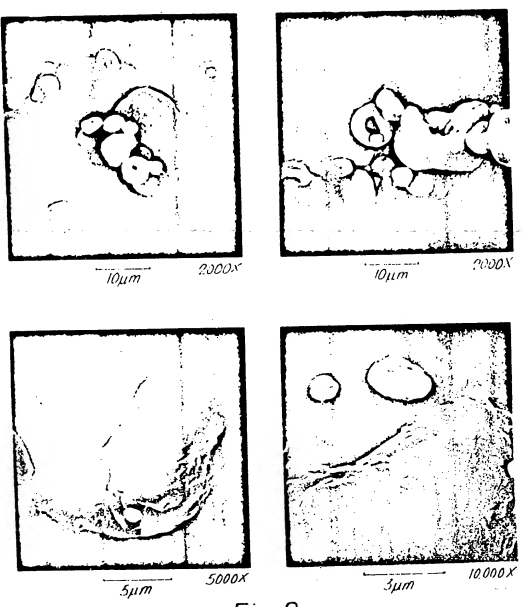


Fig.8.

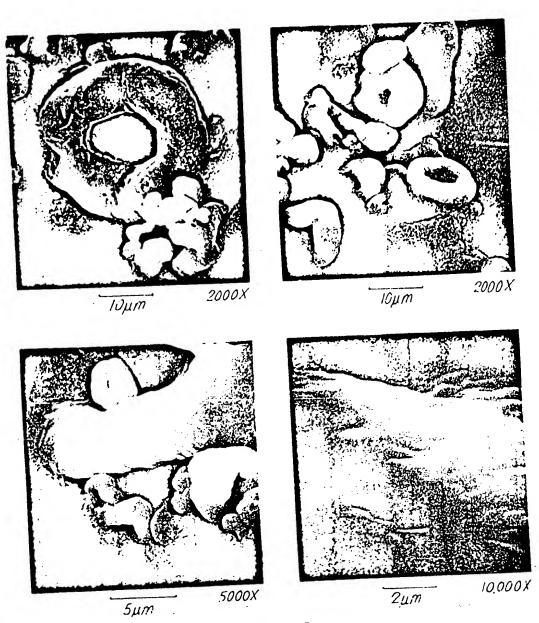
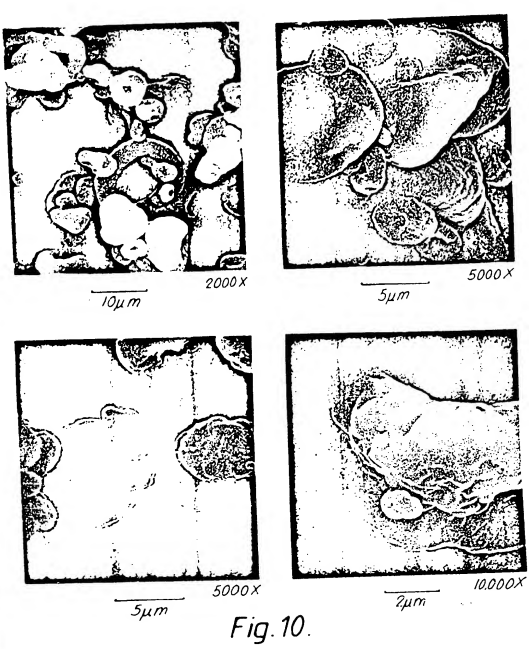


Fig.9.



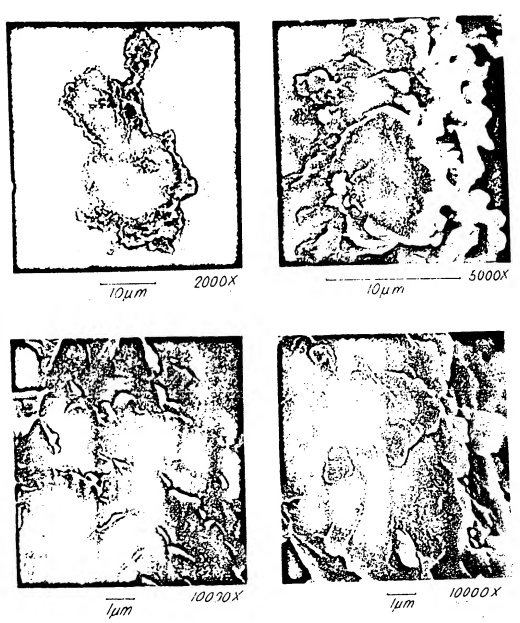


Fig.11.

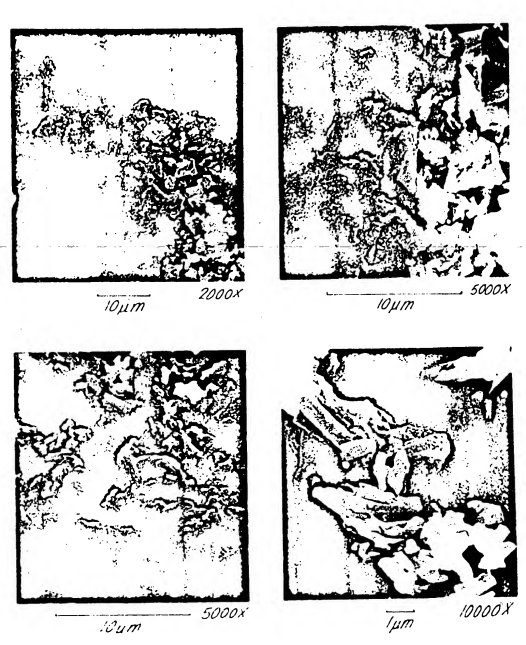


Fig.12.

SPECIFICATION

Inhalation pharmaceuticals

5. This invention relates to a novel form of drug and to methods for its production and formulation. 5 In our British Patent No. 1,122,284 we have described and claimed an insufflator device for use in the administration of powdered medicaments by inhalation. With that device, and other devices, e.g. that described in British Patent Specification No. 1,331,216, and European Patent Application No. 813021839 a user inhales air through the device which causes a powder container mounted therein to rotate. Powder 10 within the container is fluidised and dispensed into the air stream which is inhaled by the user. For optimum 10 dispensing it has been found that the powdered medicament particles should be comparatively free-flowing and yet should have an ultimate particle size of less than about ten microns to ensure adequate penetration of the medicament into the lungs of the user. These two requirements are prima facie mutually exclusive, since such fine powders are not usually sufficiently free-flowing. It has in the past been found that this 15 problem can be mitigated or overcome, e.g. as described in US Patent 4,161,516, by forming the powdered 15 medicament into small soft pellets or soft granules. Both soft pellets and soft granules will fluidise satisfactorily within the container and yet are of sufficiently low internal coherence to break up into finer particles of medicament of a therapeutically effective size in the turbulent airstream around the outside of the container. However the procedure of forming the micronised drug into soft pellets or granules is both 20 difficult and expensive. An alternative means of getting the fine particles to flow and disperse satisfactorily 20 has been to mix them with a coarse carrier, e.g. coarse lactose (see US Patent No. 3,957,965). However with all pharmaceuticals it is desirable to use as pure a form as possible (inter alia to avoid any possible adverse reactions by the patient to the excipients) and the presence of the coarse carrier is not therefore desirable. Furthermore the mixing of the fine drug with the coarse carrier involves the extra expense of the carrier, the 25 possibility of segregation of carrier and drug during transport and storage, and extra process steps which 25 add to the cost of production. Production of both the pelletised material and the blend of fine material with the coarse carrier involves the initial step of micronising the drug. Sodium cromoglycate has been made, for blending with lactose or agglomeration into soft nearly spherical pellets and administration by inhalation, as a micronised dry powder and in this form consists mostly of rods or lath-shaped crystals. In both the 30 pelletised and blended material energy is needed to break up the pellets or to separate the fine drug from the coarse carrier before or during inhalation. Thus in many instances it has also been found that the amount of drug which is available as fine particles in the air stream is dependent on the rate at which air is passed through the inhaler (i.e. the amount of energy imparted to the formulation). This can be particularly disadvantageous when the drug is used to treat patients suffering from conditions affecting their ability to 35 Thus for many years the production of drugs in a form in which they can flow easily (and therefore be filled readily into capsules) while at the same time being of a sufficiently small particle size to penetrate deep into the lung has presented a problem which has only been capable of resolution by means of complex procedures. We have now found particles which can penetrate deep into the lung and yet which are sufficiently free 40 flowing to be filled into capsules, and otherwise manipulated, without mixing with a coarse diluent or formation into soft pellets or granules. We have also found that these particles can disperse well from an inhaler at both low and high air flow rates, thus, in certain circumstances, improving consistency of capsule emptying. Furthermore we have found that the new particles can, in general, be coarser than those of the 45 prior art while giving an equivalent proportion of particles capable of penetrating deep into the lung. 45 According to the invention we provide a finely divided inhalation drug comprising a therapeutically effective proportion of individual particles capable of penetrating deep into the lung, characterised in that a bulk of the particles which is both unagglomerated and unmixed with a coarse carrier, is sufficiently free flowing to be capable of being filled into capsules on an automatic filling machine and to empty from an 50 opened capsule in an inhalation device. According to the invention we also provide a drug in finely divided and unagglomerated form, wherein a substantial proportion of the individual drug particles have a spherical, collapsed spherical, i.e. where one or both sides of the sphere appear to have been pushed inwards, or toroidal shape, i.e. the shape of a ring doughnut. The ring doughnut shapes may have a hole through the middle or may have a thin membrane 55 filling the hole. In certain cases a population of two or more of spheres, partially collapsed spheres, fully 55 collapsed spheres and ring doughnut shapes is found. The individual particles should be as rounded and smooth as possible to enable them to be carried easily In an air stream and to flow readily on capsule filling machines. We prefer the majority of the particles not to have sharp or broken edges, and for the particles themselves to be mechanically strong so that they do not

break during encapsulation or on their passage from the capsule to the lung. Thus we prefer to avoid hollow shell particles. We particularly prefer a proportion of the particles, especially when the drug is sodium cromoglycate, to be toroidal in shape. In general to shape of the particles is unrelated to particle size. We have also found that in general the particles have smooth cleavage planes, are relatively non-porous and are C'uniform density through each particle. With respect to their strength the particles of the present invention are strongly differentiated from the prior art soft pellets and granules, and with respect to their shape they

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are strongly differentiated from the prior art micronised material. A low particle density in the material is indicative of fragile particles and is, in general, to be avoided. We profer the particles to be as uniform as possible in all respects.

The surface texture of the particles will vary according to the particular drug concerned and the techniques used to produce the particles, and can vary from a highly convoluted (brain like) structure to a random fluffy or to a smooth texture. In general we prefer to avoid highly convoluted surface textures.

The roughness of the surface of the particles can be determined by measuring the total surface area of the particle by the Brunauer, Emmet and Teller (BET) method (British Standard 4359 (1969) Part 1) and comparing this with the envelope surface area of the particles as measured by permeametry (Papadnkis M. 10 (1963), Rev. Mater. Construct. Trav. 570, 79-81).

We prefer the permeametry: BET ratio to be in the range 0.5 to 1.0, preferably 0.6 to 1.0 and more preferably 0.7 to 0.97 (note a ratio of 1.0 represents a perfectly smooth particle). By way of contrast prior art micronised drugs, e.g. micronised sodium cromoglycate, have a permeametry: BET ratio of about 0.32.

We prefer the particles of the invention to be as strong and as dense as possible. The particle density of the particles (as oppossed to the bulk density) may be measured by

a) the petroleum ether method in which a known weight (25g) of powder is weighed into a measuring cylinder, a known amount of petroleum ether (50ml) is added and the mixture shaken until all the powder is suspended. The inner walls of the measuring cylinder are washed with a small amount of petroleum ether (10ml). Knowing the weight of powder used, the volume of petroleum ether added and the final suspension volume, the particle density can be calculated. or

b) the air pycnometer method in which a given amount of powder is placed in a chamber which is hermetically sealed. The volume of the chamber is gradually reduced by a moving piston until a specified pressure is reached. The position of the piston indicates the volume of the powder particles, hence the particle density can be calculated.

We prefer the particles, e.g. of sodium cromoglycate, to have a particle density according to the above methods of from about 1.3 to 1.7 and preferably from 1.3 to 1.6 g/cm³.

The micronised material, e.g. sodium cromoglycate, of the prior art has a loose bulk density of about 0.21 g/cm³ and a packed bulk density of about 0.29 g/cm³. In measuring loose bulk density a suitable amount of powder (40g) is poured, at an angle of 45°, into a measuring cylinder (250ml). The volume occupied by the powder in the measuring cylinder when related to the original mass of powder provides the measure of "loose bulk density". If the powder, in the cylinder, is tapped or jolted, e.g. using the Engelsmann Jolting Volumeter, until a stable volume is attained (500 jolts) then the lower volume after jolting when compared with the original mass of powder provides the measure of "packed bulk density".

It is also known, e.g. from British Patent Specification No. 1,549,229 that hard granules of sodium cromoglycate of particle size 60 to 200 microns (measured by sieving) can have higher bulk densities than the micronised material. However these hard granules were not designed for, and indeed would be unsuitable for, inhalation. Surprisingly we have found that the particles of the present invention have a higher bulk density than micronised material, e.g. micronised sodium cromoglycate. We prefer the particles of the present invention to have a loose bulk density of greater than about 0.3 g/cm³, preferably of greater of than 0.35 g/cm³, more preferably of from 0.35 to 0.5 g/cm³, and most preferably 0.35 to 0.4g/cm³; and a packed bulk density of from about 0.4 to 0.75 g/cm³ and preferably of from 0.55 to 0.6g/cm³. The bulk densities of materials are, in general, relative independent of the particular material used, but are dependent on the shape, size and size distribution of the particles involved.

We prefer the particles of the invention, when they comprise sodium cromoglycate and are intended for administration as a dry powder in, for example, a gelatine capsule to have a moisture content of from 5 to 14, and preferably from 8 to 11% w/w. Before filling into the capsule the powder will tend to be at the lower end of the moisture range, and after filling to be at the upper end of the range. Sodium cromoglycate powders according to the invention may also be made containing very low, e.g. less than 1%, or preferably less than 0.5%, w/w, quantities of water. These very dry powders may be used in pressurised aerosol formulations.

The water contents in this specification are those measured by drying a small sample (1 to 2g) for 15 hours at 105°C in a vacuum oven (less than 5 mm Hg) in the presence of phosphorus pentoxide.

Examples of suitable medicaments include those used for the inhalation treatment of allergic airway diseases such as pharmaceutically acceptable salts of 1,3-bis(2-carboxychromon-5-yloxy)propan-2-ol; bronchodilators, e.g. isoprenaline, salbutamol, fenoterol, terbutaline, reproterol etc and salts of anyone thereof; antibiotics, e.g. tetracycline; steroids, e.g. beclomethasone dipropionate; enzymes; vitamins and antihistamines. If desired a mixture of medicaments, for example a mixture of sodium cromoglycate and a broncholdilator, such as isoprenaline, terbutaline, fenoterol, reproterol or a salt of any one thereof, may be used. Where a highly active medicament is used which requires a small unit dose the individual particles may comprise the active ingredient together with a suitable diluent, e.g. lactose. The incorporation of the diluent in the particle avoids the possibility of segregation which is possible when individual fine particles of active ingredient are used with separate coarse particles of diluent.

We prefer that at least 50% by weight at ... preferably more than 90%, of the drug particles are of less than 60 microns, more preferably of less than 40 microns, most preferably of less than 20 microns and especially of less than 10 microns, e.g. less than 8 microns in diameter. We particularly prefer at least 50% of the particles to be of 2 to 6 microns in diameter, in general the smaller the mass mean diameter of the material

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the higher will be the dispersion of the material, as measured by the test of Example A(a).

Material according to the invention, e.g. sodium cromoglycate, having a median diameter of from 10 to 15 microns can, because of the enhanced aerodynamic properties of the particles, be equivalent in emptying and dispersion properties (see Example A) to micronised (i.e. sub 10 micron) material which has been formed into soft pellets as described in US Patent 4,161,516 or blended with coarse lactose, as described in US Patent 3,957,965.

The particle sizes in this specification are those measured with a Coulter Counter TA11 used in a standard laboratory environment, or the pipette centrifuge. In measuring particles sizes with a Coulter Counter, the sample to be analysed is dispersed in an electrolyte into which dips a glass tube. The glass tube has a 50 to 400 micron hole through the wall thereof with electrodes mounted on either side of the hole in the tube wall. The tube is immersed sufficiently for the hole and electrodes to be submerged in the liquid. The suspension is made to flow through the hole in the glass tube and as each particle passes through the orifice it displaces its own volume of electrolyte, thus changing the resistance across the hole. This change in resistance is converted into a voltage pulse with an amplitude proportional to the particle volume. The pulses are fed to an electronic counter with an adjustable threshold level such that all pulses above the threshold are counted. 15 By setting the threshold level at different values it is possible to determine the number of particles falling within given size ranges and thus the proportion of particles in a sample which fall outside a desired particle size range. The Coulter Counter measures the volume of a sphere having the same volume as the unknown material, i.e. it measures a volume diameter.

In measuring particles by the pipette centrifuge (Christison Scientific Equipment Limited) the powder is suspended in a suitable liquid (e.g. n-butanol). The suspended sample is put in a constant speed centrifuge. Samples are withdrawn from the centrifuge at selected time intervals. The level of solids in each sample is measured (normally by drying) and the average diameter calculated using an equation derived from Stokes Law (Particle Size Measurement Published by Chapman Hall 3rd Ed. Dr. T. Allen, page 377 et seq.). The 25 pipette centrifuge measures a mass, or Stokes, diameter.

The Coulter counter (with a 100 micron hole) is able to measure particle sizes of from about 2 to 40 microns

and the pipette centrifuge is able to measure particle sizes down to about 0.2 microns.

According to the invention we also provide a process for the production of finely divided drug, which comprises atomising and drying a solution of the drug and collecting some or all of the particles which are 30 below 60, preferably below 40, more preferably below 20 and especially below 10 microns in diameter. The 30 particles are preferably of the sizes given above.

Spray or flash drying of materials is well established as a drying technique in the food and other industries, but is scarcely used at all in the pharmaceutical industry. Thus spray drying is routinely used in the p. uduction of coarse particle products such as dried milk, instant coffee and dextran. The use of spray drying 35 techniques to produce very fine powders is not conventional and is unknown in the pharmaceutical field, the normal technique for producing such fine powders being to make, and then micronise, a crystalline drug. The use of a spray drying technique is advantageous in that it is adapted to suit large batch productions, thus decreasing the amount of quality control required and also in that it may remove the need for

recrystalisations and micronisation to get the drug into the desired form. Any suitable form of atomiser can be used. Atomisation results from an energy source acting on liquid bulk. Resultant forces build up to a point where liquid break-up and disintegration occurs and individual spray droplets are created. The different atomisation techniques available concern the different energy forms applied to the liquid bulk. Common to all atomisers is the use of energy to break-up liquid bulk. Centrifugal, pressure and kinetic energy are used in common forms of atomiser. Sonic and vibratory atomisers are also used. Specific atomisers which may be mentioned include rotary atomisers, e.g. those involving vaned wheels, vaneless discs, cups, bowls and plates; pressure atomisers, e.g. those involving pressure nozzles, centrifugal pressure nozzles, swirl chambers and grooved cores; kinetic energy or pneumatic atomisers, e.g. those involving two or three fluids, or internal or external mixing; and sonic energy nozzles, e.g. involving sirens or whistles. We prefer to use kinetic or pneumatic energy atomisers particularly two fluid pressure or syphon or sonic nozzle atomisers. In general two fluid pressure nozzles tend to produce powders having more desirable characteristics than two fluid syphon nozzles and two fluid

pressure nozzles also tend to give more reproducible results and use less energy. The atomiser can be used in a spray or flash drying apparatus.

The conditions of operation of the apparatus and storage of the solution (e.g. pH and temperature) should 55 clearly not be such as to degrade the drug, or introduce impurities, or biological contamination, into the

The spray drying apparatus preferably comprises the atomiser, a main chamber, one or more (e.g. two) cyclones, a bag filter and, if desired or necessary to maximise recovery, a terminal wet scrubber or electrostatic precipitator. The particle collection system is designed to capture the desired size range of particles and also to maximise the yield. All over and under size material may be recovered and recycled or put to other uses.

The solution of the drug may be in any suitable Julvent, e.g. water from a water soluble drug. The concentration of the drug in the solvent may vary over a wide range, e.g. in the case of sodium cromoglycate from 1 to 25, preferably 5 to 20, and especially 10 to 15 % w/v. In general we prefer to use a high concentration of drug as the volume and energy requirements of the atomisation and drying process are

thereby diminished. To avoid possible blockage of the atomisation device and to avoid the incorporation of unwanted impurities it is desirable to filter the solution immediately before it is passed to the atomiser. The particle size of the product tends to increase with concentration, but not rapidly, and in general concentration is not controlling with respect to particle size.

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The temperature of the air inlet and outlet to the spray drier main chamber may vary over a wide range (the range being dependent on the product being dried, the solution through put and the final moisture content required) and suitable temperatures may be found to suit each drug and solvent by simple routine experiment, in the case of aqueous solutions (of for example sodium cromoglycate), we have found that an air inlet temperature of from 180° to 350°C, preferably from 180° to 230°C, and an outlet temperature of from 10 70° to 250°C and preferably of from 70° to 120°C are suitable.

The temperature of the solution to be fed to the spray drier will vary with the drug and the solvent to be used. In general we prefer to use a temperature at which the solution can be stored for a long period in large batches without degradation. As high a temperature as possible comensurate with stability is desirable to reduce solution viscocity and provide energy to the drying process.

The air flow rate, direction into the spray drier, the temperature of the air and the rate of feed of solution to the spray drier can be optimised by simple experiment. All of the parameters in the spray drying process interrelate and can be adjusted to produce the desired product.

Gases other than air, e.g. nitrogen, can be used if desired. The use of an inert gas will be advantageous when an inflamable solvent or a readily oxidisable drug is used. The gas used, e.g. air or nitrogen, may, if 20 desired, be recycled to avoid loss of entrained drug and/or to conserve energy and the inert gas.

The particle size of the product will be set by the concentration of the feed solution, the rate of feed to the spray drier, the means of atomising the solution, e.g. the type of atomiser and the pressure of the air, and solution to be dried, the temperature and temperature gradient within the drier and, to a small extent, the air flow in the drier. The particle size and air fow will then dictate where the desired product is collected and the 25 means of collection.

The particle size of the product tends to remain fairly constant with liquid flow rate through the atomiser, but to decrease the increasing air pressure up to a limiting pressure, e.g. of about 11Kg cm⁻². The range of air pressure suitable will naturally depend on the atomisation device used, but we have found that air pressures of from about 2Kg cm⁻² to 11KG cm⁻² are in general effective, e.g. with a 0.4mm orifice syphon 30 two fluid nozzle. In order to achieve reproducible results we prefer to maintain a constant air flow to the dryer

and appropriate air flow control devices may be used if desired.

The cyclone or cyclones used to collect the dried particles are of conventional design, but adapted to collect finer particles than is normal. Thus the pressure differential across the cyclones, the combination of two or more cyclones and the design of the particular cyclones used may be adjusted to enable capture of 35 the fine particles. The bag filter used to collect the finest material is of conventional design and is readily available. The filter medium within the bag filter preferably has a high capture efficiency for particles of approximately 0.5 microns in diameter and greater. A particularly suitable medium is a polytetrafluoroethylene membrane supported on a polypropylene or polyester cloth, e.g. a needle felt cloth. Any electrostatic precipitator, or wet scrubber, used will also be of conventional design.

The product may be classified, e.g. sieved or air classified, to remove over and under sized material. The over and under sized material may be recycled or used for other purposes.

The final product may be put up in any suitable form of container such as a capsule or cartridge. Where it is desired to use the product in association with other ingredients such as colourants, sweeteners or carriers such as lactose, these other ingredients may be admixed with the particles of the invention using 45 conventional techniques or may be incorporated in the solution to be spray dried. We prefer the particles of the invention to contain medicament and water only. Mixtures of two or more different particles according to the invention, e.g. of sodium cromoglycate and a bronchodilator, such as isoprenaline sulphate or tertbutaline sulphate, may be made and filled into suitable containers.

According to our invention we also provide a method of application of a medicament, e.g. sodium 50 cromoglycate, to a patient by way of inhalation, the medicament being dispersed into an air stream, characterised in that an opened, e.g. pierced, container, e.g. capsule, containing particles according to the invention is rotated and vibrated in an air stream which is inhaled by the patient. The rotation and vibration may conveniently be produced by any one of a number of devices, e.g. the device of British Patent Specification No. 1,122,284.

The particles according to the invention may also be used in pressurised aerosol formulations (together with propellant gases, e.g. a mixture of two or more of propellants 11, 12 and 114, preferably with a surface active agent, e.g. sorbitan trioleate) or may be formed into soft pellets, e.g. as described in US Patent Specification No. 4,181,516, or may be used for application to the skin. Sodium cromoglylcate is known to be of use in the treatment of a wide variety of conditions, e.g. asthma and hay fever,

From another aspect the invention also provides a capsule, cartridge or like container containing particles according to the invention, optionally in association with other particles. We prefer the container to be loosely filled to less than about 80% by writine, preferably less than about 50% by volume, with the particles of the invention. The particles are preferably not compacted into the container. We prefer the container, e.g. capsule, to contain from 10 to 100 mg, e.g. about 20mg, of the particles. The invention will now be illustrated by the following Examples in which all perts and percentages are by

 $p(\vec{k}) \geq \min\{n, n \in \mathbb{N}\}$

	weight unless otherwise stated.	•	
٠. '	EXAMPLE 1	and the state of t	
	The active compound (A) was disso	ived in a solvent, normally water, to a concentration B (% w/v). This	_
5	solution flowed under pressure or vec	suum to the atomiser. At the atomiser the solution temperature was as of atomisation (C) and of dropist drying (D) were preset and n. The powder was captured in the drying chamber, in two cyclones	5
	March a Vancanage Buell &C 130 CV	rione of diameter 22 cm and height /4 cm and secondly a my m	
	- All -l Co-l d for l o cicloso	of diameter 14 cm) and finally in a pad liligi which had as the title.	
0	media polytetrafluoroethylene lined p vessel was weighed (E) and sized (F) (polypropylene. At the end of each run the contonts of each concerns.	10
	-2 Manufacture immediants		
	11-1 a a a a a a a a a a a a a a a a a	/v in water, and atomisation conditions (C) a pressure two fluid nozzle	
	(0.4mm orifice), a solution flow rate o	of 65ml min ⁻¹ and an atomisation pressure of 27 × 10 ³ Kg m ⁻² the	
15	results shown in Table 1 were obtained	ed.	15
	Note - Electron micrographs (see Fi	igures 1 to 4) showed.	
	Salbutamol Sulphate - smooth sph	eres	
	Terbutalene Sulphate - "orange pe	ei" spheres	
	Isoprenaline Sulphate - smooth spi	neres (3,2-g) pyran-2,8-dicarboxylic acid disodium salt "orange peel" spheres	20
20	with surface cracks	(3,2-g) pyran-2,0 dicarboxyna acta cioca	
	A M COLUMN AND N	•	
٠	Sodium Cromoglycate/)"	doughnut", apheres and other active ingredients) collapsed spheres	1
25	5 b) Varying atomisation Techniques		25
	Active ingredient (A) - Sodium Cro	moglycate.	
	Conditions used and results obtain	sed are given in Tables 2 and 28.	
	Two fluid syphon nozzie	- CT (London) Ltd. C1 Type JTA 16/50 (4min office)	
	Two fluid pressure nozzle	- CT (London) Ltd. CT Type J11	30
30	0 Ultrasonic nozzle	 Ultrasonics Ltd, 035 H Sonicore nozzle Delevan Ltd - Swirl Air Nozzle Type 32163-1 	•
	Swirl Air nozzle	- Delevan Fig - Swift Wit Mozzie 1 4be 25 100 .	
	c) Variation of powder collection tech	hniques	
	the powder is collected in the dryi	ing chamber, cyclones and a pag tilter.	35
35	Active ingredient A - Sodium Cron	noglycate.	33
	Conditions used and results obtain	ned are given in Tables 3 and 3a.	
	Powder Capture Equipment	an to (-the most is equivalent)	
	Main chamber (MC) size	- 13 cu ft (give metric equivalent) - Stairmand High Efficiency Cyclone (Diameter 14cm)	
	Cyclone A	- Vantogeren Bueil AC 130 Cyclone (Diameter 22cm, Height 74cm)	40
40		- Stairmand High Efficiency Cyclone (Diameter 11.9cm)	
	Cyclone C Bag Filter (BF)	- 1.86 M ² polytetrafluoroethylene lined polyester	
	d) Variation of droplet drying time	upon both the temperatures used in drying, i.e. air inlet temperature, the	45
4			
			1
		ASSESSED WITHIN THE INTERCONCIONES. I BUILT OF INCIDENCE WITHING TO THE PROPERTY OF THE PROPER	
	conditions used. Increased resident	ce time (i.e. slower drying) produced improved particles with improved	50
5			
	Electron micrographs of a selection	on of the above powders are shown in the accompanying Figures. Figure	1

11 and 12 are electron micrographs of, respectively polletised sodium cromoglycate, and micronised sodiu cromoglycate and are included for comparison purposes only. In each of Figures 1 to 12 the magnification and an approximate scale is given.

TABLE 1

Electron Micrograph Figure No		1 (B cyclone)			2 (B cyclone)	м		•	
Powder Recovered E/F Cyclone Cyclone B A micron volume median diameter	18/3.4	17/4.0	1.202	3333	15/4.1	173.6	25/4.2	123.2	937.8 (cyclone C)
Powder Ra Cyclone B micron	807.5	83/4.3	78/4.1	34/8.5	786.2	756.6	587.4	757.0	
Main Chamber	2.0.	÷	÷	Ŕ	7/16.5	à	-/21-	13/19.0	<i>‡</i>
D) Air Flow Rate m³s-1	0.034	9:03	0.034	0.034	0.034	0.034	0.034	0.034	9.034
Drying Conditions (D) Inlet Outlet Temp. Temp.	8	102	501	8	8	101	88	90	.100
Drying Inlet Temp.	2 6	202	\$	5	50	500	220	502	500
Active Ingredient (A)	Sodium Cromoglycate	Terbutalene Sulphate	Salbutamol Sulphate	Boprenaline Sulphate	4,6-Dioxo-10-propyl-4H,8H-pyrano[3,2-g] pyran-2,8-dicarboxylic acid disodium salt	Sodium Cromoglycate (100)/ Terbutalene Sulphate (0.522) w/w	Sodium Cromoglycate (100)/ Sulbutamol Sulphate (0.522) w/w	Sodium Cromoglycate (100)/ Isoprensline Sulphate (0.522) w/w	Sulbutamol Sulphate* (1.8)/ Lactose (100) w/w
<u> </u>	_	<u>د</u> ؛	است		ú	ಕ	۲.	ದ	ශ්

*Cyclone configuration changed to MCC/BF.

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ž ž	Run Atomiser No. Type	Atomisati Solution Conc	Atomisation Conditions (C) Solution Solution Ato Conc Feed Rate Pre: % w/v Ls -1x10-3 Kgr	(C) Atomisation Pressure Kgm ⁻ ×10³	Drying Co Inlet Temp. °C	Drying Conditions (D) Inlet Outlet Temp. Temp.	Air Flow Rate m³s-1	Power Rec Main Chamber	Power Recovered Erf Main Cyclone Chamber B micron median	Cyclone A volume diameter	Bed Fifter	Electron Micrograph Figura No	
ಠ	Slotted Disc	5	0.57	23000	220	136	9.034	•	91/15	95.2		5 (B cyclone)	
=	Slowed Disc	9	0. 48	Ę	214	8	0.034	Ŕ	78/22	2/4.0		•	
: :	Teles	2	0.70		230	318	0.034	Ŕ	71/39	34.3			
2 :		5	0.50		215	127	0.034	21/24	7.71/87			6 (B cyclone)	
<u>≠</u>	Two Fluid Syphon	ıo	0.33	150.7	538	125	0.034	1/-	19/4.5	31,2.8	.		
مو	Nozzłe	8	1.33	150.7	82	3	0.034	26/15.5	127.4	62/3.1			
, ≝		5	0.90	56.4	210	90	0.034	<i>*</i>	708.5	23/3/0			
17.		5	0.හ	105.7	225	113	90.03	ኤ	344.7	31/2.9	302.1		•
<u> </u>	Two Fluid	51	0.37	28.2	8	132	9.034	823	62/6.8	30.3.7			
<u> </u>		9	0.33	28.2	900	8	9.03	12.	77/9.2	11/3.5			•
Ŕ		5	1.52	18.3	210	፯	0.034	24.	74/16.0	2/4.0		5	-
*		5	0.42	39.5	203	137	0.034	525	53/10	33.4	9.30	7 (A cyclone)	
ä		5	1.33	36.6	5 2	88	0.034	7 ,	77/10.5	103.2			
ដ	Pressure Nozzle 5mm orifice	6	1.17	21.1	302	8	0.034	. 12.	79/9.2	9,4.2			
*	. Ultresonic Nozzłe	5	1.47	35.2	210	83	0.034	ઢં	82/9.6	123.3			
*	26. Swirt Air	15	1.17	49.3	200	8	0.034	13.	79/14.5		ക്	ത	
ž	ejzze	E C	ber contents sl	•Chamber contents showed incomplete drying	te drying.			. '					

M.C.) size Air Pycn- Petro Loose Pecked ometer Ether Ether Awww volume diameter 15 2.6 15 2.7 1.35 1.45 0.42 0.58 7.0 2.1 1.35 1.45 0.43 0.63 1.00 4.3 1.77 1.58 1.68 0.34 0.48 8.5 2.4 1.33 1.45 1.59 1.68 0.34 0.48 8.5 2.4 1.33 1.45 1.59 1.59 1.59 1.59 1.59 1.59 1.59 1.5	2 4	Dispersion (see Exam-	Coulter	Particle Density g/cm³		Bulk Density g/cm³	usity	Moisture	Emptying (see Example Ab)	BET	Peramea- metry	Permeatmery BET ratio
worker median diameter	G.	() 4	ę.	Air Pycn- ometer		Loose	Packed	<u>;</u>	3	Ę. B	× 10	
2.6 15 86 68 1.4 5.2 1.35 1.45 0.42 0.58 7.0 0.62 0.486 0 1.7 1.7 1.56 1.45 0.74 5.5 88 0.48 0.33 8.6 4.3 1.56 1.56 0.74 5.5 88 0.48 0.33 8.6 15.5 1.59 1.69 0.34 0.48 8.5 59.2 2.42 1.25 2.4 1.33 1.45 1.5 1.55 0.31 0.43 0.43 1.75 1.1 2.4 1.33 1.45 1.5 0.31 0.43 1.5 96 1.75 1.1 2.4 8.5 8.5 8.5 8.5 1.75 1.1 1.1 1.1 1.1 1.1 1.1 1.1 1.2 1.2 1.2 1.2 1.2 1.2 1.2 1.2 1.2 1.2 1.2 1.2 1.2 1.2	•	****	volume median diameter					} *	Ř			
1.4 5.2 1.35 1.45 0.42 0.58 7.0 0.62 0.486 0 1.7 1.7 1.56 1.45 0.43 0.63 25 0.48 0.33 10.0 4.3 1.56 0.50 0.74 5.5 88 0.48 0.33 11.7 1.56 1.50 0.74 5.5 88 0.48 0.33 2.9 1.55 1.66 0.34 0.48 8.5 57 1.55 2.4 1.33 1.45 1.69 0.34 0.48 8.5 592 1.75 1.1 2.6.1 4.2 1.55 0.31 0.43 28 1.75 1.1 2.4 3.5 1.45 1.5 0.31 0.43 28 1.75 1.1 2.4 3.5 3.5 3.5 3.5 1.75 1.1 2.4 3.5 3.5 3.5 3.5 1.7 1.1 2.4		12.6	5						&	•		
22 1.35 1.45 0.42 0.58 7.0 0.62 0.496 0 177 0.43 0.63 88 0.48 0.33		. 717	. 52		•		•	•	8		•	•
17 0.43 0.63 88 55 10.0 4.3 5.5 88 0.48 0.33 17.7 1.56 0.50 0.74 5.5 88 0.48 0.33 8.6 1.53 <td>.</td> <td></td> <td>R</td> <td>1.35</td> <td>1.45</td> <td>0.42</td> <td>0.58</td> <td>7.0</td> <td></td> <td>0.62</td> <td>964.0</td> <td>6.0 E</td>	.		R	1.35	1.45	0.42	0.58	7.0		0.62	964.0	6.0 E
40.0 4.3 5.5 88 0.48 0.33 17.7 1.56 0.50 0.74 5.5 88 0.48 0.33 8.6 15.9 1.56 0.34 0.48 8.5 59.2 2.42 1.25 21.4 2.8 1.59 1.66 0.34 0.48 8.5 59.2 2.42 1.25 18.6 9.2 1.55 1.55 0.31 0.43 28 1.75 1.1 24.1 1.56 1.55 0.31 0.43 28 1.75 1.1 12.3 14.5 1.55 0.31 0.43 28 1.75 1.1 24.4 9.5 1.55 0.31 0.43 28 1.75 1.1 24.4 9.5 1.55 0.31 0.43 28 1.75 1.1 24.4 9.5 1.55 0.31 0.43 28 1.75 1.1 94.5 9.5 9.63 <		•	17			0.43	9.63		88	•	•	
17.7 1.56 0.50 0.74 5.5 88 0.48 0.33 8.6 1.59 1.56 0.34 0.48 8.5 57 7 21.4 2.8 1.59 1.66 0.34 0.48 8.5 59.2 2.42 1.25 18.6 9.2 1.55 0.31 0.43 28 1.75 1.1 12.3 14.5 1.56 1.55 0.31 0.43 28 1.75 1.1 24.4 9.5 1.56 1.55 0.31 0.43 28 1.75 1.1		0.0 4	4.3	•			•	•	ĸ		•	
2.9	<u>ન</u>		17.7	1.56	•	0.50	0.74	5.5	&	0. 48	8 .0	2
8.6 15.6 1.68 0.34 0.48 8.5 59.2 2.42 1.25 21.4 2.8 1.69 1.60 0.34 0.48 8.5 59.2 1.25 18.6 9.2 1.33 1.45 1.5 1.5 0.31 0.43 2.8 1.75 1.1 24.1 4.2 1.56 1.55 0.31 0.43 2.8 1.75 1.1 24.4 9.5 1.55 0.31 0.43 2.8 1.75 1.1	#	•	2.9				•		23	•	•	•
21.4 2.8 1.59 1.66 0.34 0.48 8.5 59.2 2.42 1.25 19.6 9.2 9.2 9.2 1.56 1.55 0.31 0.43 28 1.75 1.1 24.1 4.2 1.56 1.55 0.31 0.43 28 1.75 1.1 24.4 9.5 9.6 96.3 96.3 96 96	卓		15.5	•	4		•	•	8 -			
24 1.33 1.45 98 19.6 9.2 92 26.1 4.2 1.56 0.31 0.43 28 1.75 1.1 12.3 14.5 6.9 96.3 7 1.4 24.4 9.5 96 96	7.	21.4	2.8	1.59	8 .	0.34	0.48	8.5	59.2	2.42	क्	7 6
19.6 9.2 28 1.75 1.1 26.1 4.2 1.56 1.55 0.31 0.43 28 1.75 1.1 12.3 14.5 6.9 96.3 24.4 9.5	Ŕ	•	24	1.33	1.45	•			8			•
26.1 4.2 1.56 1.55 0.31 0.43 28 1.75 1.1 12.3 14.5 6.9 96.3 24.4 9.5	Z,		9.2	•	•	•	•		8	. !	. :	
12.3 14.5 . 6.9	ង		7	35.	35 .		0.43		8	6.1 6.1	3	}
24.4 9.5	7		14.5	•	•	•	•	63	96.3		•	•
	Ŕ		9.5	•	٠.		•		8	•	• .	•

		Atomisation Conditions (C)	nditions	(C)	E	Dryir In is	Drying Conditions (D)	g Conditions (D) Outlet Air Flow	Main	Powder R	Powder Recovered (E/F) Cyclone Cyclone C	JF) Cyclone Bag	₽
돌호	Run Powder No. Capture Exclorent	Atomiser		SolutionSolution Conc. Feed Rate	sation Pressure	Temp.	Temp.	Rate	Chamber	< <	8	υ Έ	Filter
	Configuration	Type	\$ \$	Ls-1×10-3	Kgm ⁻² ×10 ³	ပ္	۰۲	m3s-1		median	diameter	microns	
Ŕ	MCABBF	Two Fluid Syphon Nozzle	6	1.17	105.7	210	88	0.034	Ř	87/9.6	10/4.2		
Ŕ	MCBF	Nozzłe	5	1.27	105.7	215	88	0.034	14/17				865.2
, ह		Nozzłe	9	98.0	105.7	218	112	0.034	Ŕ	402.9	35.6.4	~	272.0
Ŕ		Two Fluid Pressure	5	č:	18.3	8	8	0.034				LO.	50/13.5
Ŕ	MC8F	Nozzłe 4mm	5	0.42	8.58	8	120	0.034	•223			on .	96/5.2
3.	MCBABE	Orifice	5	1.52	18.3	210	호	0.034	24 /-	3/4.0	37.6		
8	MCCBF		6	6.0	35.2	2 6	88	0.034	11/-			86/6.5	તું.
ន់	, MCBF	Two Fluid Pressure	0	1.73	16.2	185	74	0.034	-/19		Š	e	39/14
ಕ	MCB/ABF	Nozzie Smm	5	1.16	21.1	5 6	8	0.034	1 5	94.2	7881	711/38	
8	36. MC/CBF	Orifice	15	1.23	26.8	222	102	0.034	. 5			}	

c ,	Run Dispersion No. (see Exam- ple Ac)	Pipette Centrifuge perticle	Coulter particle size	Particle g/cm³	Density	Bulk Density g/cm³	nsity	Moisture	Moisture Emptying (see Example Ab)	ᇤ	Peramea- metry	Peramea-Permeametry metry BET ratio
•		e s		Air Pycn- ometer	Petro- leum Ether	Loose	Loose Packed	3	*	m²kg ⁻¹ ×10³	95	
•-	******	mass median diameter	volume median diameter				٠		:	· ·		
	25.4	•	42		•				.			
	8.3	•	17.0		•		•		8			
•		1.7	2.0	•	•	•		•	8			
-	17.1	•	13.5		•			•	93			
-			24.0	1.33	1.45	•			88			•
	20.6		89		•				8			
ផ	20.0		14.0		•		•		91	•		
	19.6 26.1		9.2	1.56	. 4.55	0.31	. 0.43		8 8	1.75	1.12	. 35.
×	8.05	•	11.5	•	•	•		28	92.9	•	•	•

CABLE 4

	Electron Micrograph Figure	•			10 (1st cyclone)		
	Air Flow Rate	- 5 E	0.034	9.034	9.03		0.034
Drying Conditions	Outlet Temp.	ပ္	88	*	122		8
Drying C		မှ	<u> </u>	345	8		140
	Atomisation Pressure	0 ⁻³ Kgm ⁻² ×10 ³	176.2	55.0	35.2		23.3
Atomisation Conditions	Solution Feed Rate	Ls ⁻¹ ×10 ⁻³	1.67	0.48	0.67		1.28
Atomisation	Solution Conc.	% %/	8	vs	01		5
	Atomiser Type		Two Fluid	Syphon Nozzłe	Two Fluid	Pressure Nozzie 4mm	Orifice
	æ & •		*	3.	×		R i

EXAMPLE 2

EXAMPLE 2

The experiment was carried out using a spray drier which had a main chamber and a single cyclone. (Main chamber 0.37m³, cyclone Stairmand High Efficiency design with diameter 119mm). Atomisation was achieved using a two fluid pressure nozzle with orifice diameter 0.44mm. With an aqueous sodium cromoglycate feed solution concentration of 15 % w/v, an air flow rate of 0.034M³s⁻¹ and other conditions set out in Table 5, the results shown in Tables 5, 5s and 5b v.sre obtained. Table 5b gives test results when the powders produced according to this Example have been filled into hard gelatine capsules.

				TABLE 5	·			*	10
	Atomisation Co	nditions (C)	Dry	ing Cond	itions (D)	Powder f	Recovered E	/F `	• •
Run	Solution Feed	Atomisation	n Inle	t	Outlet	Main	Cyc	lone	
No.	Rate	Pressure	Ten	np	Temp	Chambe	r		15
	Ls ⁻¹ ×10 ⁻³	Kgm ⁻² ×10	•с		℃			ledian	
40.	1.33	27.5	190	-200	70-80	33/-	67/	13.0	20
41.	1.58	21.2	220)-230	85-95	40/-	60/	14.7	
42.	1.43	25.4	195	5-200	80-90	20/-	80	13.8	25
43 .	1.50	24.0	199	5-204	75-85	33/-	67	/13.7	
44.	1.58	22.6	190	0-200	70-80	36/-	64	/14.0	
45.	1.50	24.0	19	5-205	80-90	34/-	66	/16.5	30
				TABLE	5a				
					data .				35
Te	est		40 40	41	42	43	44	45	
M	loisture % w/w		8.8	9.7	8.4	9.8	, 9.8	9.5	40
		neter microns	13.0	14.7	13.8	13.7	14.0	16.5	
9%	w/w 6 microns		10	8	9	8	8	7	
*	w/w 30 microns		4	7	. 8	8	, в	15	45
L	oose Bulk Density	g/cm³	0.39	0.38	0.39	0.38	0.36	0.37	
F	acked Bulk Densit	ly g)cm³	0.58	0.56	0.58	0.57	0.57	0.59	
	No. 40. 41. 42. 43. 44. 45.	Run Solution Feed No. Rate Ls ⁻¹ ×10 ⁻³ 40. 1.33 41. 1.58 42. 1.43 43. 1.50 44. 1.58 45. 1.50 Test Moisture % w/w Particle Siza: Volume median dian % w/w 6 microns % w/w 30 microns Loose Bulk Density	No. Rate Pressure Ls ⁻¹ ×10 ⁻³ Kgm ⁻² ×10 ⁻³ 40. 1.33 27.5 41. 1.58 21.2 42. 1.43 25.4 43. 1.50 24.0 44. 1.58 22.6 45. 1.50 24.0 Test Moisture % w/w Particle Size: Volume median diameter microns % w/w 6 microns	Atomisation Conditions (C) Dry Run Solution Feed Atomisation Inle No. Rate Pressure Ten La⁻¹×10⁻³ Kgm⁻²×10³ °C 40. 1.33 27.5 190 41. 1.58 21.2 220 42. 1.43 25.4 190 43. 1.50 24.0 190 44. 1.58 22.6 190 45. 1.50 24.0 190 Test Run Nut 40 Moisture % w/w 8.8 Particle Siza: Volume median diameter microns 13.0 % w/w 6 microns 10 % w/w 30 microns 4 Loose Bulk Density g/cm³ 0.39	Atomisation Conditions (C) Run Solution Feed Atomisation Inlet No. Rate Pressure Temp Ls -1 × 10 -3 Kgm -2 × 10 3 °C 40. 1.33 27.5 190-200 41. 1.58 21.2 220-230 42. 1.43 25.4 195-200 43. 1.50 24.0 196-204 44. 1.58 22.6 190-200 45. 1.50 24.0 195-205 TABLE Solution Feed Atomisation Inlet Moisture % w/w 8.8 9.7 Particle Size: Volume median diameter microns 13.0 14.7 % w/w 6 microns 10 8 % w/w 30 microns 10 8 % w/w 30 microns 4 7 Loose Bulk Density g/cm² 0.39 0.38	Run Solution Feed Atomisation Inlet Outlet No. Rate Pressure Temp Temp La ⁻¹ ×10 ⁻³ Kgm ⁻² ×10 ³ *C *C 40. 1.33 27.5 190-200 70-80 41. 1.58 21.2 220-230 85-95 42. 1.43 25.4 195-200 80-90 43. 1.50 24.0 195-204 75-85 44. 1.58 22.6 190-200 70-80 45. 1.50 24.0 195-205 80-90 TABLE 5a Powder data Run Number 40 41 42 Moisture % w/w 8.8 9.7 8.4 Particle Siza: Volume median diameter microns 13.0 14.7 13.8 % w/w 6 microns 10 8 9 %w/w 30 microns 4 7 8 Loose Bulk Density g/cm² 0.39 0.38	Atomisation Conditions (C)	Run Solution Feed Atomisation Inlet Outlet Main Cyc No. Rate Pressure Temp Temp Chamber Ls⁻¹×10⁻³ Kgm⁻²×10³ °C °C %/Micron Volume M Diameter 40. 1.33 27.5 190-200 70-80 33/- 67/ 41. 1.58 21.2 220-230 85-95 40/- 80/- 42. 1.43 25.4 195-200 80-90 20/- 80/- 43. 1.50 24.0 195-204 75-85 33/- 67/- 44. 1.58 22.6 190-200 70-80 36/- 64 45. 1.50 24.0 195-205 80-90 34/- 66 TABLE 5a Powder data Moisture % w/w 8.8 9.7 8.4 9.8 9.8 Particle Size: Volume median diameter microns 10.8 9 8 8 % w/w 6 microns<	Run Solution Feed Atomisation Inlet Outlet Main Cyclone No. Rate Pressure Temp Temp Chamber La⁻¹×10⁻³ Kgm⁻²×10³ °C °C %/Micron Volume Median Diameter 40. 1.33 27.5 190-200 70-80 33/- 67/13.0 41. 1.58 21.2 220-230 85-95 40/- 60/14.7 42. 1.43 25.4 195-200 80-90 20/- 80/13.8 43. 1.50 24.0 195-204 75-85 33/- 67/13.7 44. 1.58 22.6 190-200 70-80 36/- 64/14.0 45. 1.50 24.0 195-205 80-90 34/- 68/16.5 TABLE 5a Powder data Run Number 40 41 42 43 44 45 Moisture % w/w 8.8 9.7 8.4 9.8 9.8 9.

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	Test	<i>Caj</i> Run Numb	osule Data					
5	1631	40	41	42	43	44	45	5
	Moisture Content % w/w							
	Powder when in the capsule	12.1	11.9	12.2	12.2	13.3	13.2	10
10	Capsule shell	13.9	14.2	13.3	13.5	13.1	13.0	10
	Total mg/capsule	11.8	11.9	11.9	11.6	11.6	11.5	
15	Emptying Test % w/w							15
	<i>(See Example Ab)</i> Mean	95.4	96.4	97.1	97.2	97.4	96.2	
	Range	87.3-99.1	92.6-99.3	93.1-100	95.5-98.9	92.7-100	94.3-98.2	20
20	Dispersion mg/capsule (See Example Ac)	5.32	4.03	4.74	4.97	4.28	3.12	20

EXAMPLE 3

25 Pressure nozzle

The trial was carried out using a spray drier having a main chamber and a single cyclone.

This experiment was used to demonstrate that the pressure nozzle was capable of providing small particles and establishing the order of magnitude of pressure required to produce particles with an average mass mean diameter of less than 10 microns. An atomiser pressure of 2.1×106 Kgm⁻², a feed concentration 30 of 6% w/v of aqueous sodium cromoglycate, an air inlet temperature of 230°C and an air outlet temperature of 120°C was used. The resulting powder had particles of size 11 microns mass mean diameter with a particle bulk density similar to that of micronised powder, but with a tapped bulk density twice that of micronised powder. The powder was satisfactory in the capsule emptying test.

The appearance of the powder under the light microscope was of uniform spheres or collapsed spheres 35 with negligible fractured particles.

EXAMPLE A

The drug is dispensed from a gelatine capsule 6.4 mm in diameter and having two holes 0.8mm in diameter in a shoulder thereof mounted in a device (commercially available under the Registered Trade Mark 'Spinhaler') according to British Patent No. 1,122,284 having a drawn wire shaft 2.03mm diameter journalied in a hard nylon bearing tube 13mm long and having an internal diameter of 2.08mm at its inner end (i.e. that end housing the free end of the shaft) and of 2.44mm at its other end.

The particles are preferably such that when put up in gelatine capsules 6.4mm in diameter each containing 20mg of the particles they meet the criteria set out in the tests below:

(a) Dispersion test

The filled capsules are mounted in the capsule holder of the powder insufflator (having the specific dimensions set out immediately above) of British Patent Specification No. 1,122,284 and pierced to produce two holes of 0.8mm diameter in a shoulder of the capsule. The dispersion of the medicament in the cloud 50 delivered by the insuffiator is determined using a modified version of the multistage liquid impinger described in British Patent Specification No. 1,081,881. The modifications incorporated in the present design are the addition of an extra impingement stage, and of a glass tube with a right angled bend approximately mid-way along its length. The extra impingement stage was added prior to the three stages described in British Patent Specification No. 1,081,881 and consists essentially of a jet of internal diameter 2.5cm and a 56 collection plate of diameter 5cm designed to give an effective cut-off of approximately 12 microns at an air flow rate of 80 litres per minute. The glass tube, also of internal diameter 2.5cm abutts the external end of the jet of the extra stage. The insuffiator is inserted into the upper, horizontal end of the glass tube and air drawn through at 60 litres per minute for 30 seconds. At least five capsules are treated in this manner and the results are averaged. The weight of the medicament collected on each stage of the impinger, on the glass tube, and on a filter paper positioned after the final stage is determined spectrophotometrically after solution 60 in an appropriate volume of distilled water (or by any other appropriate method).

The particles disperse satisfactorily if an averr - stotal for each capsule of at least 0.5 mg, preferably at least 2.timg and most proferably at least 5.0mg of the particles are found on a combination of the last two stages and filter paper of the multi-stage liquid impinger.

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(b) Emptying test

The filled capsules are mounted in the capsule holder of the powder insufflator (having the specific dimensions set out above) of British Patent Specification No. 1,122,284 and pierced to produce two holes of 0.8mm diameter in a shoulder of the capsule. The insufflator is placed in a device adapted to suck air through 5 It for 2.5 seconds, the air flow rate at no time exceeding 60 litres per minute, and being held at 60 litres per minute for at least 2 seconds. The capsule mounted in the insufflator is subjected to 4 sucks as described and the weight of the material remaining in the capsule is determined. The above procedure is repeated 20 times

and the average of the results determined. The capsules empty satisfactorily if an average of at least 50%, preferably at least 75% and most preferably 10 at least 90% by weight of the material has emptied from each capsule.

(c) Dispersion

In a further refinement, the multistage liquid impinger of Example Aa) was simplified to give a single stage 15 liquid impinger, consisting of a single impingement assembly with a filter downstream. The impingement assembly consisted of a vertical jet of internal diameter 1.9cm and a collection plate of diameter 3.8cm. At the upper end, the jet was bent through an angle of 90° and the insufflator was attached to the distal end of this horizontal portion. The impingement characteristics of this single stage device were intended to be such that material reaching the filter of this device is similar in particle size to that reaching the final two stages 20 and filter of the multistage liquid impinger of Example Aa). The percentage of material reaching the filter of

the device is determined. In all samples of sodium cromoglycate prepared by the techniques exemplified above at least some of the particles were of toroidal (ring doughnut) shape.

25 CLAIMS

- 1. A finely divided inhalation drug comprising a therapeutically effective proportion of individual particles capable of penetrating deep into the lung, characterised in that a bulk of the particles which is both unagglomerated and unmixed with a coarse carrier, is sufficiently free flowing to be filled into capsules on 30 an automatic filling machine and to empty from an opened capsule in an inhalation device.
 - 2. An inhalation drug in finely divided and unagglomerated form, wherein a substantial proportion of the individual drug particles have a spherical, collapsed spherical or ring doughnut shape.
 - 3. A drug according to Claim 2 which contains sodium cromoglycate and wherein the particles are of ring doughnut shape.
- 4. A finely divided inhalation drug, wherein the permeametry: BET ratio, as hereinbefore defined, is in the range 0.5 to 1.0. 5. A drug according to Claim 4, wherein the ratio is from 0.6 to 1.0.
 - A drug according to Claim 5, wherein the ratio is from 0.7 to 0.97.
 - A drug according to any one of the preceding claims, wherein the particle density is from 1.3 to 1.7 g 6.
- 8. A drug according to Claim 7, wherein the particle density is from 1.3 to 1.6 g/cm³. 40 cm³.
 - A drug according to any one of the preceding claims, having a loose bulk density of greater than 0.3g/cm³.
 - 10. A drug according to Claim 9 having a loose bulk density of greater than 0.35g/cm³.
 - 11. A drug according to Claim 10 having a loose bulk density of from 0.35 to 0.5g/cm³. A drug according to any one of the preceding claims having a packed bulk density of from 0.4 to 12.
- 13. A drug comprising sodium cromoglycate, wherein more than 90% of the drug particles are less than 0.75g/cm³. 60 microns in diameter and the drug has a loose bulk density of greater than 0.3g/cm³.
- 14. A drug comprising sodium cromoglycate, wherein more than 90% of the drug particles as less than 60 microns in diameter and the drug has a packed bulk density of from 0.4 to 0.75g/cm³
 - 15. A drug according to any one of the preceding claims, which is sodium cromoglycate and contains from 5 to 14% w/w of water.
 - 16. A drug according to Claim 15, which contains from 8 to 11% w/w water.
- 17. A drug according to any one of Claims 1 to 14, which is sodium cromoglycate and contains less than 1% w/w of water.
 - 18. A drug according to Claim 17, which contains less than 0.5% w/w of water. A drug according to any one of the preceding claims which comprises a mixture of sodium
- cromoglycate and a bronchodilator. 20. A drug according to any one of the preceding claims, wherein at least 50% of the drug particles are
 - 21. A drug according to Claim 20, whr _in at least 50% of the drug particles are less than 40 microns in less than 60 microns in diameter.
- 22. A drug according to Claim 21, wherein at least 50 % of the drug particles are less than 20 microns in diameter.

diameter.

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1 to 3.

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42. A drug produced by a process according to any one of Claims 33 to 41.

41. A process according to Claim 33 and substantially as hereinbefore described in any one of Examples

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